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<p>2 DETERMINATION OF SPECIES ORIGIN</p> <p>2.1 Quality Control of Antiserums</p> <p>2.1.1 Before using any new lot number of precipitating antiserum for testing casework samples, the specificity must be tested and appropriately documented in the laboratory's quality control (QC) records. Routine quality control testing will be performed at the time the antiserum is reconstituted, or if the antiserum is received in liquid form, within one week of receipt.</p> <p>2.1.2 Anti-human serum, not anti-human hemoglobin, will be used with the procedure in this section for determining whether a sample is of human origin. Anti-human serum, as well as all animal antisera in the "Routine Species Collection" specified below, must be tested against all available species (normal or whole sera and known bloods) in the "Routine Species Collection".</p> <p>2.1.3 A positive control, a host control (typically normal rabbit serum or normal goat serum), and a negative control (distilled water) must be included in the specificity testing. The host control (representing the animal in which the antiserum was prepared) is used to demonstrate that the antiserum is not reacting with any proteins in the animal in which it was made.</p> <p>2.1.4 The quality control documentation will include:</p> <p>2.1.4.1 A diagrammatic representation of the placement of samples in the wells of the rosette on the Ouchterlony plate with the subsequent observations (i.e., precipitin lines).</p> <p>2.1.4.2 Date of the testing.</p> <p>2.1.4.3 Initials of the person conducting the testing.</p> <p>2.1.4.4 Lot number, date of receipt, and manufacturer of the antiserum being tested.</p> <p>2.1.4.5 Lot number, date of receipt, and manufacturer of the normal sera being used for the testing.</p> <p>2.1.4.6 Results of the testing.</p> <p>2.1.5 Once the appropriate testing has been performed on a particular lot number of antiserum, it need not be repeated for each case. If another vial of the same lot number is received on a different date, the QC testing described above must be repeated.</p> <p>2.2 Quality Control of Normal (Whole) Sera</p> <p>2.2.1 Before using any new lot number of normal serum for testing casework samples, the specificity must be tested and appropriately documented in the laboratory's quality control (QC) records. Routine quality control testing will be performed at the time the normal serum is reconstituted, or if the normal serum is received in liquid form, within one week of receipt.</p> <p>2.2.2 All normal sera in the "Routine Species Collection" must be tested against all available antisera in the "Routine Species Collection".</p>	

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<p>2.2.3 A positive control must be included in the specificity testing. Although normal human serum may be purchased, a <u>straw colored dilution</u> of known human blood may be used instead. Similarly, the use of known blood from other species may replace the purchase and use of normal serums from the species.</p> <p>2.2.4 It is not necessary to conduct quality control testing on the known bloods. Spot the liquid blood sample onto a stain card and label known bloods with the species name and date of preparation/initials of person preparing the sample. Store the dried known bloods in the freezer.</p> <p>2.2.5 The quality control documentation will include:</p> <p>2.2.5.1 A diagrammatic representation of the placement of samples in the wells of the rosette on the Ouchterlony plate with the subsequent observations (i.e., precipitin lines).</p> <p>2.2.5.2 Date of the testing.</p> <p>2.2.5.3 Initials of the person conducting the testing.</p> <p>2.2.5.4 Lot number, date of receipt, and manufacturer of the normal serum being tested.</p> <p>2.2.5.5 Lot number, date of receipt, and manufacturer of the antisera being used for the testing.</p> <p>2.2.5.6 Results of the testing.</p> <p>2.2.6 Once the appropriate testing has been performed on a particular lot number of normal serum, it need not be repeated for each case. If another vial of the same lot number is received on a different date, the QC testing described above must be repeated.</p> <p>2.3 Routine Species Collection</p> <p>2.3.1 The following will be maintained in each laboratory as the “Routine Species Collection” and will undergo QC testing for specificity as outlined above:</p> <p>2.3.1.1 Bovine antiserum and normal bovine serum or known blood</p> <p>2.3.1.2 Swine antiserum and normal swine serum or known blood</p> <p>2.3.1.3 Cat antiserum and normal cat serum or known blood</p> <p>2.3.1.4 Dog antiserum and normal dog serum or known blood</p> <p>2.3.1.5 Rabbit antiserum and normal rabbit serum or known blood</p> <p>2.3.1.6 Sheep antiserum and normal sheep serum or known blood</p> <p>2.3.1.7 Deer antiserum and normal deer serum or known blood</p> <p>2.3.1.8 Human antiserum and normal human serum or known blood</p> <p>2.3.1.9 Normal goat serum or known blood (goat antiserum is unavailable)</p>	

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<p><b>NOTE:</b> It is recommended that the antisera and normal sera listed above be reconstituted and QC tested at the time of receipt to ensure ready availability.</p> <p>2.3.2 Other antisera/normal sera, such as bear, rodent, fowl, horse, etc. should also be maintained for use in special cases and must also undergo QC testing as specified above. However, since these are used only in special cases, it is recommended that they not be reconstituted, aliquoted, and QC tested until it is determined that there is a specific need to do so.</p> <p>2.4 Storage of Antiserum/Normal Serum</p> <p>2.4.1 Small aliquots of the antiserum/normal serum will be prepared for routine use and frozen within one week of reconstitution (when antiserum is lyophilized) or upon receipt (when antiserum is liquid). All frozen aliquots have an indefinite expiration date.</p> <p>2.4.2 A thawed aliquot may be stored refrigerated for up to 1 month. If the aliquot is to be maintained in this manner, the expiration date must be clearly marked on the vial. Otherwise, the aliquot will be immediately discarded following its use.</p> <p>2.5 Labeling of Antiserum/Normal Serum</p> <p>2.5.1 Labels on each aliquot will include:</p> <p>2.5.1.1 The manufacturer</p> <p>2.5.1.2 Type of antiserum or normal serum</p> <p>2.5.1.3 Lot number</p> <p>2.5.1.4 Date reconstituted/date frozen</p> <p>2.5.1.5 Initials of the person preparing the aliquot</p> <p>2.6 OUCHTERLONY (DOUBLE DIFFUSION) TEST (Reference 5, pp. 221-241, Appendix A)</p> <p>2.6.1 Equipment</p> <p>2.6.1.1 Punch</p> <p>2.6.1.2 Aspirator</p> <p>2.6.1.3 100 ml and 500 ml graduated cylinders</p> <p>2.6.1.4 Balance</p> <p>2.6.1.5 Spatula</p> <p>2.6.1.6 Scissors</p> <p>2.6.1.7 Tweezers</p>	

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<p>2.6.1.8 Hot plat or oven (37° C)</p> <p>2.6.1.9 Incubator (optional)</p> <p>2.6.1.10 Magnetic stir plate</p> <p>2.6.1.11 Refrigerator (optional)</p> <p>2.6.2 Materials</p> <p>2.6.2.1 Petri dishes, slides, or comparable containers</p> <p>2.6.2.2 Test tubes</p> <p>2.6.2.3 Weigh boat or weigh paper</p> <p>2.6.2.4 Moisture chamber</p> <p>2.6.2.5 Disposable pipets</p> <p>2.6.2.6 Capillary tubes</p> <p>2.6.3 Reagents</p> <p>2.6.3.1 Normal saline (0.9%)</p> <p>2.6.3.2 Agarose gel (1%)</p> <p>2.6.3.3 Distilled water</p> <p>2.6.3.4 Antiserum</p> <p>2.6.3.5 Normal serum or known blood (positive control)</p> <p>2.6.4 Agarose Gel Preparation</p> <p>2.6.4.1 Normal saline (0.9% NaCl):</p> <ul style="list-style-type: none"> <li>• 9 g Sodium chloride</li> <li>• 1000 ml Distilled water</li> <li>• Mix thoroughly until dissolved.</li> </ul> <p>2.6.4.2 Agarose gel (1%):</p> <ul style="list-style-type: none"> <li>• 1 g Type I agarose</li> <li>• 100 ml Normal saline (0.9% NaCl)</li> <li>• Heat until agarose is dissolved. Allow to cool slightly.</li> </ul> <p>2.6.4.3 Pour the agarose into a petri dish, onto a slide, or into a comparable container to a thickness of 2-3 mm and allow to cool.</p>	

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<p>2.6.4.4 Cut wells in a rosette pattern (refer to the diagram below) into the gel using a punch or disposable pipet connected to an aspirator.</p> <div> <p>The diagram shows a large circle representing a gel plate. Inside this circle, there are seven smaller circles arranged in a rosette pattern. One circle is in the center, and six circles are arranged around it in a hexagonal pattern. The peripheral wells are numbered 1 through 6, starting from the top right and moving clockwise. The central well is not numbered.</p> </div> <p>2.6.5 Storage and Labeling</p> <p>2.6.5.1 When a batch of plates is prepared, the plates should be numbered consecutively and placed in a moisture chamber in the refrigerator. Label the moisture chamber with the lot number of the batch (date of preparation/initials of person preparing the plates).</p> <p>2.6.5.2 There is no expiration date (see 2.6.6 Minimum Standards and Controls).</p> <p>2.6.6 Minimum Standards and Controls</p> <p>2.6.6.1 A positive control (known sample against which the antiserum is directed) and a negative control (distilled water) will be tested on each plate. A substrate control may be tested when appropriate.</p> <p>2.6.7 OUCHTERLONY DOUBLE DIFFUSION PROCEDURE</p> <p>2.6.7.1 To prepare an extract of the stain, place a small cutting of the stain in distilled water or saline until a <u>straw color</u> is obtained. A small piece of stained material, which is moistened with distilled water or saline, can be used in lieu of an extract. If a substrate control is utilized treat this sample in the same manner as the stain.</p> <p>2.6.7.2 Add antiserum in the center well of the Ouchterlony plate with a disposable pipet or capillary tube.</p> <p>2.6.7.3 Add appropriate extracts/pieces of stained material, the positive control, and negative control to the surrounding wells. Do not overfill the wells. Avoid getting bubbles in the wells. Document the placement of samples in the wells of the rosette on the Ouchterlony work sheet found in Appendix B.</p>	

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<p>2.6.7.4 Record the lot numbers of antisera and normal sera used for the testing procedure.</p> <p><b>NOTE:</b> Alternatively, known normal serum/known blood extract may be placed in the center well with appropriate antisera in the surrounding wells or the stain extract/piece of stained material may be placed in the center well with appropriate antisera in the surrounding wells.</p> <p>2.6.7.5 Incubate the plate in a moisture chamber at 37° C for 3-4 hours. Alternatively, it may be left overnight at room temperature or 4° C.</p> <p>2.6.7.6 Record observations (precipitin lines) on the diagram, and interpret and record the results.</p> <p>2.6.7.7 All controls must give the expected results before a conclusion can be reached on an unknown sample, i.e., white precipitin lines must be observed between the antiserum and positive control (known serum) and no precipitin lines should be observed between the antiserum and the negative control.</p> <p>2.6.7.8 Interpretation</p> <p>2.6.7.8.1 Positive Result = White precipitin lines between the antiserum well and the sample well</p> <p>2.6.7.8.2 Negative Result = No precipitin lines between the antiserum well and the sample well</p> <p>2.6.7.8.3 Inconclusive Result = Questionable precipitin lines between the antiserum well and the sample well, OR precipitin lines between the antiserum well and the negative control well(s), OR no precipitin line between the antiserum well and the positive control well. If sufficient sample remains, an inconclusive result should be repeated.</p> <p><b>NOTE:</b> The prozone phenomenon can result in a soluble antigen-antibody complex due to too many antibodies present to form a complete lattice (Reference 6, Appendix A). Because of this phenomenon, weak precipitin lines may be observed initially, but disappear upon staining with Coomassie Brilliant Blue R250. This is considered an inconclusive result. Therefore, it is recommended that the testing results be recorded PRIOR TO staining as well as after staining.</p> <p>2.6.7.9 Staining the plate with Coomassie Brilliant Blue R250 may be necessary to visualize weak reactions. Refer to 2.7 for the Coomassie Brilliant Blue R250 staining procedure.</p> <p>2.6.7.10 Reporting Results</p> <p>2.6.7.10.1 Report positive test results as “<u>(species tested according to the label on the antiserum)</u> protein was identified...”</p> <p>2.6.7.10.2 Report negative test results as “no <u>(species tested according to the label on the antiserum)</u> protein was identified...”</p> <p>2.6.7.10.3 Report inconclusive test results as “the test for <u>(species tested according to the label on the bottle)</u> protein was inconclusive...”</p>	

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2.7 COOMASSIE BRILLIANT BLUE R250 STAINING PROCEDURE (Reference 6, Appendix A)	
2.7.1 Safety Considerations	
2.7.1.1 Coomassie Brilliant Blue R250 - Caution! Avoid contact and inhalation! Emits toxic fumes under fire conditions! Container explosion may occur under fire conditions!	
2.7.1.2 Methanol - Caution! Irritant! Dangerous when exposed to heat or flame!	
2.7.1.3 Glacial acetic acid - Caution! Corrosive! Flammable!	
2.7.2 Equipment	
2.7.2.1 Weight	
2.7.2.2 Oven or incubator (40-60°C)	
2.7.2.3 Rotator (optional)	
2.7.2.4 10 ml, 50 ml, and 500 ml graduated cylinders	
2.7.2.5 Balance	
2.7.2.6 Spatula	
2.7.2.7 Trays for staining and destaining	
2.7.3 Materials	
2.7.3.1 Gel bond, glass plate, or other support medium	
2.7.3.2 Weigh boats or weigh paper	
2.7.3.3 Whatman #1 filter paper	
2.7.3.4 Paper towels	
2.7.4 Reagents	
2.7.4.1 Staining Solution	
2.7.4.2 Destaining Solution	
2.7.4.3 Normal saline (0.9% NaCl) – refer to 2.7.8.1	
2.7.4.4 Distilled water	

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<p>2.7.5 Preparation of Staining and Destaining Solutions</p> <p>2.7.5.1 Staining Solution</p> <ul style="list-style-type: none"> <li>• 0.1 g Coomassie Brilliant Blue R250</li> <li>• 45.0 ml Methanol</li> <li>• 10.0 ml Glacial acetic acid</li> <li>• 45.0 ml Distilled water</li> <li>• Mix the above ingredients until thoroughly dissolved.</li> </ul> <p>2.7.5.2 Destaining Solution</p> <ul style="list-style-type: none"> <li>• 45.0 ml Methanol</li> <li>• 10.0 ml Glacial acetic acid</li> <li>• 45.0 ml Distilled water</li> <li>• Mix the above ingredients until thoroughly dissolved.</li> </ul> <p>2.7.6 Storage</p> <p>2.7.6.1 The Staining and Destaining Solutions are stable at room temperature.</p> <p>2.7.7 Labeling</p> <p>2.7.7.1 Label as Staining or Destaining Solution with the lot number (date of preparation followed by the initials of the person preparing the solution).</p> <p>2.7.7.2 There is no expiration date.</p> <p>2.7.8 COOMASSIE BRILLIANT BLUE R250 STAINING PROCEDURE</p> <p>2.7.8.1 Wash the plate for two hours to overnight in normal saline solution (0.9% NaCl - 9 g NaCl in 1000 ml distilled water) to remove unprecipitated proteins.</p> <p>2.7.8.1.1 If pieces of stained material were used in lieu of extracts, remove these prior to washing the plate.</p> <p>2.7.8.1.2 The gel may detach from the plate during the washing process. Mark the orientation of the gel to ensure that it can be re-oriented properly after the washing has been completed.</p> <p>2.7.8.1.3 Alternatively, the gel may be removed from the plate before beginning the washing process. If this is done, mark the gel to ensure that it can be re-oriented properly after the washing has been completed.</p> <p>2.7.8.2 The next day wash the gel for approximately fifteen minutes in distilled water. Rinse the gel and repeat the wash. Ensuring proper orientation, place the gel (face up) on the hydrophilic side of a piece of gel bond or on some other support medium such as a glass plate.</p>	



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<div> 2.7.8.3 Cover the gel with a piece of Whatman #1 filter paper moistened with distilled water. Add a layer of paper towels on top of the filter paper and press with a weight for approximately 30 minutes. Remove paper towels and filter paper and dry the gel in a 40-60° C oven. </div> <div> 2.7.8.4 Place the gel in the staining solution and allow it to soak for less than 1 minute up to 10 minutes. This may be done on a rotator. Intermittently check staining progress to prevent over staining. </div> <div> 2.7.8.5 Place the gel in the destaining solution until the background is clear or until no more dye leaches from the gel. This may be done on a rotator. Change the destaining solution and destain further if desired. </div> <div> 2.7.8.6 Record observations (precipitin lines) on the diagram, and interpret and record the results. All controls must give the expected results before a conclusion can be reached on an unknown sample, i.e., blue precipitin lines must be observed between the antiserum and positive control (known serum) and no precipitin lines should be observed between the antiserum and the negative control. </div> <div> <div>2.7.8.7 Interpretation</div> <div> 2.7.8.7.1 Positive Result = Blue precipitin lines between the antiserum well and the sample well. </div> <div> 2.7.8.7.2 Negative Result = No precipitin lines between the antiserum well and the sample well. </div> <div> 2.7.8.7.3 Inconclusive Result = Questionable precipitin lines between the antiserum well and the sample well, OR precipitin lines between the antiserum well and the negative control well(s), OR no precipitin line observed between the antiserum well and the positive control well. If sufficient sample remains, an inconclusive result should be repeated. </div> </div> <div> <div>2.7.8.8 Reporting Results</div> <div>2.7.8.8.1 Refer to 2.6.7.10.</div> </div>	

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